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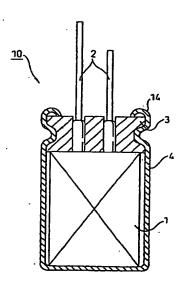
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(54) ELECTROLYTE FOR DRIVING ELECTROLYTIC CAPACITOR AND ELECTROLYTIC CAPACITOR

(57)An electrolytic solution for use in an electrolytic capacitor, comprising a solution containing a solvent consisting of 20 to 80% by weight of an organic solvent and 80 to 20% by weight of water, and at least one electrolyte selected from the group consisting of a carboxylic acid or a salt thereof and an inorganic acid or a salt thereof, having added thereto at least one nitro compound selected from the group consisting of nitrophenitrobenzoic acid, dinitrobenzoic nitroacetophenone and nitroanisole. The electrolytic solution has a low impedance and excellent low-temper-, ature stability, along with good working life characteristics, and it can also exhibit an excellent hydrogen gas absorption function when an electrolytic solution contains a highly increased amount of water in its mixed solvent or when an electrolytic capacitor is used under high temperature conditions.





Description

Technical Filed

[0001] The present invention relates to an electrolytic capacitor. More particularly, the present invention relates to an electrolytic solution for use in an electrolytic capacitor, which has a low impedance and excellent low-temperature stability, along with good characteristics of working life, and an electrolytic capacitor using the same, specially an aluminum electrolytic capacitor.

10 Background Art

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[0002] Generally, a capacitor is a general electrical part and is widely used for as a power supply circuit, a noise filter and a digital circuit component in various electric/electronic parts. Capacitors are roughly classified into electrolytic capacitors and other capacitors such as ceramic capacitors, film capacitors, etc.

[0003] Various types of electrolytic capacitors are used at present and examples thereof include aluminum electrolytic capacitors, wet tantalum electrolytic capacitors and the like. It is an aluminum electrolytic capacitor from which a particularly excellent function is expected in the present invention. Therefore, the present invention will now be described with reference to this kind of an electrolytic capacitor. The term "electrolytic capacitor" used herein refers to an aluminum electrolytic capacitor unless otherwise stated.

[0004] A conventional aluminum electrolytic capacitor can be produced typically by using an anode foil, which is made by etching a high-purity aluminum foil to thereby increase its surface area, and anodizing the surface of the aluminum foil to provide an oxide film, and a cathode foil whose surface has only been etched. The resulting anode foil and cathode foil are disposed opposite each other and an element with a wound structure is made by interposing a separator (release paper) between those foils and then the element is impregnated with an electrolytic solution. The element impregnated with the electrolytic solution is contained in a case (generally made of aluminum), which is then sealed with an elastic sealant, thus completing an electrolytic capacitor. Electrolytic capacitors also include electrolytic capacitors other than those with a wound structure.

[0005] In the above-described electrolytic capacitor, the characteristics of the electrolytic solution may be a large factor which decides the performance of the electrolytic capacitor. With the size reduction of the electrolytic capacitor, an anode foil or cathode foil having a large surface area produced by etching has been used and the resistivity of the capacitor has recently increased. Therefore, an electrolytic solution having a low resistivity (specific resistance) and thus high conductivity is required as an electrolytic solution to be used in the electrolytic capacitor.

[0006] A conventional electrolytic solution for use in an electrolytic capacitor is generally prepared by dissolving, as an electrolyte, a carboxylic acid such as adipic acid, benzoic acid, etc. or an ammonium salt thereof into a solvent prepared by adding about 10% by weight or less of water to ethylene glycol (EG) as a principal solvent. Such an electrolytic solution has a specific resistance of about $1.5 \Omega \cdot m$ ($150 \Omega \cdot cm$).

[0007] On the other hand, the capacitor is required to have a low impedance (Z) to sufficiently exert the performance thereof. The impedance is decided by various factors and, for example, it is reduced when the electrode area of the capacitor increases. Therefore, an attempt to reduce the impedance is made as a matter of course in case of a large-sized capacitor. An attempt to reduce the impedance by improving a separator has also been made. However, the specific resistance of the electrolytic solution is a large controlling factor, particularly in a small-sized capacitor.

[0008] A lower-specific resistance electrolytic solution using an aprotic organic solvent such as GBL (γ -butyrolactone) has recently been developed (see, Japanese Unexamined Patent Publication (Kokai) Nos. 62-145713, 62-145714 and 62-145715). However, the capacitor using this aprotic electrolytic solution is by far inferior in impedance to a solid capacitor using an electronic conductor having a specific resistance of 1.0 Ω • cm or less.

[0009] The aluminum electrolytic capacitor has poor low-temperature stability because of use of an electrolytic solution, and a ratio of an impedance at -40°C to that at 20°C (100 kHz), Z (-40°C)/Z (20°C), is as large as about 40 at present. Under these circumstances, it is now required to provide an aluminum electrolytic capacitor which has a low impedance and excellent low-temperature stability.

[0010] Further, water used as portion of the solvent in the electrolytic solution of the aluminum electrolytic capacitor is a chemically active substance to aluminum constituting the anode foil or cathode foil. Accordingly, there is a problem that water reacts with the anode foil or cathode foil, thereby to generate a hydrogen gas and to drastically deteriorate the performance as a capacitor.

[0011] To solve a problem such as generation of hydrogen gas found in a load life test of the electrolytic capacitor, a trial of absorbing the generated hydrogen gas has hitherto been made. For example, Japanese Examined Patent Publication (Kokoku) No. 59-15374 discloses an electrolytic solution, for use in operation of an electrolytic capacitor, produced by adding a carboxylic acid and an ammonium salt of the carboxylic acid to a solvent having added thereto 5 to 20% by weight of water, thereby to prepare a buffer solution and further adding 0.05 to 3% by weight of p-nitrophenol

to the buffer solution. When using this electrolytic solution, there can be provided an electrolytic capacitor wherein low-temperature stability and a working life characteristics are improved by inhibiting the occurrence of the boehmite reaction and generation of the hydrogen gas.

[0012] Japanese Unexamined Patent Publication (Kokai) No. 63-14862 also discloses an electrolytic solution for use in the operation of an electrolytic capacitor capable of exhibiting an excellent corrosion preventing function against washing with a halogenated hydrocarbon, which is produced by adding o-nitroanisole to an electrolytic solution prepared by dissolving various organic acids, inorganic acids or salts thereof in a solvent composed exclusively of ethylene glycol. This publication describes that o-nitroanisole used as a corrosion inhibitor has a hydrogen gas absorption function, that is, a function of absorbing a hydrogen gas generated from the interior during the use of the electrolytic capacitor, thereby making it possible to inhibit an accident of safety-vent operation and a change in capacitance.

[0013] However, it has been found, as a result of the present inventors' study, that p-nitrophenol or o-nitroanisole can exhibit an initial hydrogen absorption function in the case of a conventionally used electrolytic solution of low water concentration for use in operation of an electrolytic capacitor, but cannot exhibit and maintain a satisfactory hydrogen gas absorption function when a content of water is 20% by weight or more based on the solvent in the electrolytic solution or when the electrolytic capacitor is operated under high temperature conditions for a long period of time.

Disclosure of the Invention

[0014] The present invention has been accomplished to solve the above problems of the prior art, and an object thereof is to provide an electrolytic solution, for use in an electrolytic capacitor, which has a low impedance and excellent low-temperature stability, expressed by an a ratio of an impedance at low temperature to that at normal temperature, along with good characteristics of working life, and also it can exhibit an excellent hydrogen gas absorption function even when an electrolytic solution contains a highly increase amount of water in its mixed solvent or when an electrolytic capacitor is used under high temperature conditions.

[0015] Another object of the present invention is to provide an electrolytic capacitor using the electrolytic solution of the present invention, specially an aluminum electrolytic capacitor.

[0016] These objects as well as other objects of the present invention will easily become apparent from the following detailed description.

[0017] In one aspect thereof, the present invention resides in an electrolytic solution for use in an electrolytic capacitor, comprising a solution containing a solvent consisting of 20 to 80% by weight of an organic solvent and 80 to 20% by weight of water, and at least one electrolyte selected from the group consisting of a carboxylic acid or a salt thereof and an inorganic acid or a salt thereof, having added thereto at least one nitro compound selected from the group consisting of nitrophenol, nitrobenzoic acid, dinitrobenzoic acid, nitroacetophenone and nitroanisole.

[0018] In the electrolytic solution of the present invention, the nitro compound can exhibit an excellent hydrogen absorption function in combination with the other electrolytic solution component on even when the nitro compound is used alone. To obtain a more remarkable function, two or more nitro compounds are used in combination, more preferably.

[0019] When the nitro compound is added to the electrolytic solution of the present invention, the nitro compound is added in the amount of 0.01 to 5% by weight based on the total amount of the electrolytic solution.

[0020] The organic solvent to be used, along with water, to form a mixed solvent is a protic solvent, an aprotic solvent, or a mixture thereof. That is, the protic solvents and aprotic solvents may be used alone or two or more kinds of them may be optionally used in combination, respectively. The protic solvent is preferably an alcohol compound, while the aprotic solvent is preferably a lactone compound.

[0021] The carboxylic acid or salt thereof to be used as the electrolyte in the electrolytic solution of the present invention is preferably at least one selected from the group consisting of formic acid, acetic acid, propionic acid, butyric acid, p-nitrobenzoic acid, salicylic acid, benzoic acid, oxalic acid, malonic acid, succinic acid, glutaric acid, adipic acid, fumaric acid, maleic acid, phthalic acid, azelaic acid, citric acid and hydroxybutyric acid, and ammonium, sodium, potassium, amine and alkyl ammonium salts thereof.

[0022] The inorganic acid or salt thereof which is also used as the electrolyte is at least one selected from the group consisting of phosphoric acid, phosophorous acid, hydrophosphorous acid, boric acid, sulfamic acid, and ammonium, sodium, potassium, amine and alkyl ammonium salts thereof.

[0023] In addition to the nitro compound, additives selected from the group consisting of the following group:

- (1) a chelate compound,
- (2) saccharides,
- (3) hydroxybenzyl alcohol and/or L-glutamic-diacetic acid or a salt thereof, and
- (4) gluconic acid and/or gluconic lactone may be optionally contained in the electrolyte of the present invention. These additives may be used alone, or two or more kinds of them may be optionally used in combination.

[0024] In another aspect thereof, the present invention resides in an electrolytic capacitor comprising an electrolytic solution for use in an electrolytic capacitor which comprises a solution containing a solvent consisting of 20 to 80% by weight of an organic solvent and 80 to 20% by weight of water, and at least one electrolyte selected from the group consisting of a carboxylic acid or a salt thereof and an inorganic acid or a salt thereof, having added thereto at least one nitro compound selected from the group consisting of nitrophenol, nitrobenzoic acid, dinitrobenzoic acid, nitroacetophenone and nitroanisole.

Brief Description of the Drawings

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Figure 1 is a sectional view showing one preferred embodiment of the electrolytic capacitor according to the present invention, and

Fig. 2 is a perspective view showing the constitution of a capacitor element of the electrolytic capacitor shown in Fig. 1.

Best Mode for Carrying Out the Invention

[0026] As described above, the electrolytic solution for an electrolytic capacitor according to the present invention is characterized by containing:

at least one nitro compound selected from the group consisting of nitrophenol, nitrobenzoic acid, dinitrobenzoic acid, nitroacetophenone and nitroanisole,

in addition to a solvent consisting of 20 to 80% by weight of an organic solvent and 80 to 20% by weight of water, and

at least one electrolyte selected from the group consisting of a carboxylic acid or a salt thereof and an inorganic acid or a salt thereof.

[0027] In the electrolytic solution for an electrolytic capacitor according to the present invention, a solvent containing a highly increased amount of water, which consists of a mixture of an organic solvent and water, is used as the solvent for dissolving the electrolyte.

[0028] As described above, protic solvents or aprotic solvents are used alone or, optionally, in combination. Examples of preferred protic solvent include alcohol compound. Specific examples of the alcohol compound used advantageously herein include, but are not limited to, monohydric alcohol such as ethyl alcohol, propyl alcohol, and butyl alcohol; dihydric alcohol such as ethylene glycol, diethylene glycol, triethylene glycol, and propylene glycol; and trihydric alcohol such as glycerin. Examples of preferred aprotic solvent include lactone compounds. Specific examples of the lactone compounds used advantageously herein include, but are not limited to, γ -butyrolactone and other intramolecular polarizable compounds. When using at least one solvent selected from the protic and aprotic solvents in the practice of the present invention, more specifically, one protic solvent may be used, one aprotic solvent may be used, plural protic solvents may be used, plural aprotic solvents may be used, alternatively a mixed solvent of at least one protic solvent and at least one aprotic solvent may be used.

[0029] In the electrolytic solution of the present invention, water is added in addition to the above-described organic solvents as the solvent component. Particularly, the present invention differs from a conventional electrolytic solution in that a comparatively large amount of water is used. According to the present invention, by using such a solvent, the solidifying point of the solvent is lowered, thereby making it possible to improve the specific resistance at low temperature of the electrolytic solution and to realize good low-temperature stability, expressed by a ratio of a resistivity at low temperature to that at normal temperature. A content of water in the electrolytic solution is preferably within a range from 20 to 80% by weight, and an organic solvent is contained as a balance. When the content of water is smaller than 20% by weight and when the content of water exceeds 80% by weight, the degree of depression in solidifying point of the electrolytic solution becomes insufficient, thereby making it difficult to obtain good low-temperature stability of the electrolytic capacitor. A preferred content of water in the solvent is within a range from 30 to 80% by weight, and a most preferred content of water in the solvent is within a range from 45 to 80% by weight.

[0030] As the electrolyte in the electrolytic solution of the present invention, an organic acid, particularly preferably a carboxylic acid or a salt thereof, and an inorganic acid or a salt thereof may be used. These electrolyte components may be used alone, or two or more kinds of them may be used in combination.

[0031] Examples of carboxylic acid which can be used as the electrolyte component include, but are not limited to, monocarboxylic acid such as formic acid, acetic acid, propionic acid, butyric acid, p-nitrobenzoic acid, salicylic acid, and benzoic acid; and dicarboxylic acid such as oxalic acid, malonic acid, succinic acid, glutaric acid, adipic acid, fumaric

acid, maleic acid, phthalic acid, and azelaic acid. Carboxylic acids having a functional group such as hydroxyl group, for example, citric acid and hydroxybutyric acid may also be used.

[0032] Examples of the inorganic acid which can also be used as the electrolyte component include, but are not limited to, phosphoric acid, phosphorous acid, hypophosphorous acid, boric acid and sulfamic acid.

[0033] As the salt of the above-described carboxylic acid or inorganic acid, various salts can be used. Preferred salts include, for example, ammonium salts, sodium salts, potassium salts, amine salts and alkyl ammonium salts. Among these salts, an ammonium salt is preferably used.

[0034] In addition, using the inorganic acid or salt thereof as the electrolyte in the practice of the present invention, depression in solidifying point of the electrolytic solution can be expected, thereby making it possible to contribute to a further improvement in low-temperature stability of the electrolytic solution. The use of the inorganic acid or salt thereof is noticeable in that the hydrogen gas absorbability (described in detail hereinafter) derived from the nitro compound used particularly in the present invention can be maintained for a long period of time.

[0035] According to the present inventors' study, by using an electrolyte such as inorganic acid or salt thereof in combination with the above-described carboxylic acid or salt thereof, an effect of remarkably prolonging a working life of the electrolytic capacitor, occurs, as compared with the case where they are used alone. Furthermore, an inorganic acid-based electrolyte has hitherto been used exclusively in a medium to high-voltage (160 to 500 volts) type electrolytic capacitor in a conventional electrolytic capacitor in view of the conductivity. However, when using electrolytes in combination, like the present invention, the electrolyte can also be used advantageously in a low-voltage (lower than 160 volt) type electrolytic capacitor.

[0036] The amount of the electrolyte used in the electrolytic solution of the present invention can be appropriately determined depending on various factors such as characteristics required to the electrolytic solution and the capacitor obtained finally, kind, compositions and amount of the solvent, and kind of the electrolyte. As described above, when using the inorganic acid-based electrolyte in combination with the carboxylic acid-based electrolyte, the amount of the inorganic acid-based electrolyte contained in the mixed electrolyte can vary within a wide range, but the inorganic acid-based electrolyte is preferably contained in the amount within a range from about 0.1 to 15%, by weight, based on the total amount of the electrolyte.

[0037] The electrolytic solution of the present invention is characterized by further adding, as an additive, at least one nitro compound selected from the group consisting of a nitrophenol such as p-nitrophenol, a nitrophenol acid such as p-nitrophenol acid, a dinitrophenol acid, a nitrophenone such as p-nitrophenone and nitrophenole, to an electrolytic solution of the above-described specific compositions, that is, an electrolytic solution comprising an aqueous mixed solvent consisting of 20 to 80% by weight of an organic solvent and 80 to 20% by weight of water, and at least one electrolyte selected from the group consisting of a carboxylic acid or a salt thereof and an inorganic acid or a salt thereof.

[0038] In the present invention, a particularly hydrogen gas absorption function could be confirmed when using the above-described group of nitro compounds, but the actual reasons have not yet been clarified. However, it is considered, based on the present inventors' experience, that a large factor is in that substituents contained in each nitro compound exhibit the hydrogen gas absorption function at different timings. The nitro compound used herein can also have a function of inhibiting corrosion of the element caused by a function of a halogenated hydrocarbon used on washing of a printed circuit board, for example, trichloroethane (a halogen capturing function, in other words).

[0039] When the nitro compound is added to the electrolytic solution of the present invention, the nitro compound can exhibit satisfactory hydrogen gas absorption functions and halogen capturing functions even when used alone because specific compositions effective for the function of the present invention are employed in the electrolytic solution itself. According to the present inventors' finding, a more preferred function can be expected from using two or more nitro compounds in combination. It is generally recommended to use two nitro compounds in combination. The nitro compound is preferably added in the amount within a range from 0.01 to 5% by weight based on the total amount of the electrolytic solution. When the amount of the nitro compound is smaller than 0.01% by weight, an expected function is hardly obtained. On the other hand, even when the amount exceeds 5% by weight, a further improvement in expected function cannot be expected and a deleterious influence is sometimes exerted on the other characteristics.

[0040] The use of the nitro compound will be described in more detail below. The absorption function of the hydrogen gas generated on the reaction between aluminum and water is liable to be lowered with the increase in amount of water in the solvent when using the nitro compound alone, as was described in the item entitled "Background Art". This lowering tendency becomes drastic in the case where the electrolytic solution is subjected to high temperature conditions. However, problems caused by using these nitro compounds alone can be solved by using two or more nitro compounds in combination, as in the present invention. Actually, in case of the electrolytic solution of the present invention, the hydrogen gas absorbability can be maintained under high temperature conditions for a longer period of time, than in the case where these nitro compounds are used alone, by using plural nitro compounds.

[0041] An excellent function in absorption of the hydrogen gas according to the present invention could also be confirmed in a relation to the electrolyte used in combination. In a conventional electrolytic solution, the procedure of adding

only one nitro compound to only a carboxylic acid-based electrolyte, or adding only one nitro compound to only an inorganic acid-based electrolyte has been employed. However, a satisfactory hydrogen gas absorption function cannot obtained by the procedure in case where the amount of water contained in the solvent is large, and the same results are obtained in an electrolytic solution wherein both of a carboxylic acid-based electrolyte and an inorganic acid-based electrolyte are present. In case of the electrolytic solution of the present invention (using only one nitro compound), the hydrogen gas absorbability could be, surprisingly, maintained for a longer period of time than the case where nitro compounds are used alone even in case of the carboxylic acid/inorganic acid mixed electrolytic solution.

[0042] The electrolytic solution of the present invention can optionally contain, as an additive, components other than those described above. Preferred additives include, for example, the following compounds.

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- (1) Chelate compound, for example, ethylenediamine-N,N,N',N'-tetraacetic acid (EDTA), trans-1,2-diaminocy-clohexane-N,N,N',N'-tetraacetic acid, monohydrate (CyDTA), N,N-bis(2-hydroxyethyl)glycine (DHEG), ethylenediamine-N,N,N',N'-tetrakis(methylenephosphonic acid) (EDTPO), diethylenetriamine-N,N,N',N'',N''-pentaacetic acid (DTPA), 1,3-diamino-2-hydroxypropane-N,N,N',N'-tetraacetic acid (DPTA-OH), ethylenediamine-N,N'-diacetic acid (EDDA), ethylenediamine-N,N'-bis(methylenephosphonic acid), hemihydrate (EDDPO), O,O'-bis(2-aminoethyl)ethyleneglycol-N,N,N',N'-tetraacetic acid (EDTA), N-(2-hydroxyethyl)ethylenediamine-N,N',N'-triacetic acid (EDTA-OH) and others. The chelate compound is preferably added in the amount within a range from 0.01 to 3% by weight. Such a chelate compound can exert effects such as prolongation of a working life due to inhibition of the hydration reaction of an aluminum (Al) electrode foil of a low-impedance capacitor, improvement in low-temperature stability of an electrolytic capacitor (a change between an impedance at normal temperature and that at low temperature decreases because the solvent has compositions close to those corresponding to a non-frozen state), and improvement in corrosion resistance.
- (2) Saccharides, for example, glucose, fructose, xylose, galactose and others. The saccharides are preferably added in the amount within a range from 0.01 to 5% by weight. These saccharides can exert effects such as prolongation of a working life due to inhibition of the hydration reaction of an aluminum electrode foil of a low-impedance capacitor, inhibition of decomposition or activation of an electrolyte (e.g. carboxylic acid) due to the addition of saccharides, and improvement in low-temperature stability of an electrolytic capacitor (a change between an impedance at normal temperature and that at low temperature decreases because the solvent has compositions close to those corresponding to a non-frozen state).
- (3) Hydroxybenzyl alcohol, for example, 2-hydroxybenzyl alcohol, L-glutamic-diacetic acid or a salt thereof and others. This additive is preferably added in the amount within a range from 0.01 to 5% by weight. Such an additive can exert effects such as prolongation of a working life due to inhibition of the hydration reaction of an aluminum electrode foil of a low-impedance capacitor, and improvement in low-temperature stability of an electrolytic capacitor (a change between an impedance at normal temperature and that at low temperature decreases because the solvent has compositions close to those corresponding to a non-frozen state).

The above-described compounds (1) to (3) can exhibit various remarkable effects by adding them to the electrolytic solution of the present invention, and almost all of the effects can be expected even in case where no nitro compound is contained in the electrolytic solution. According to the present inventors' study, these excellent effects can be obtained by using at least one of the above compounds (1) to (3) in combination with gluconic acid or gluconic lactone described below.

In addition to the above-described additives (also including the case where nitro compounds are added alone), the electrolytic solution of the present invention can optionally contain:

- (4) gluconic acid and gluconic lactone alone or in combination. This kind of the additive is preferably added in the amount within a range from 0.01 to 5% by weight. Gluconic acid and gluconic lactone can further exert remarkable effects such as improvement in corrosion resistance, in addition to functions, which are specific to the present invention, such as prolongation of a working life of an electrolytic capacitor, improvement in low-temperature stability and excellent hydrogen gas absorption function, by containing them in the electrolytic solution of the present invention.
- [0043] In addition to the above-described additives, additives conventionally used in the field of aluminum electrolytic capacitors and other electrolytic capacitors may also be added. Preferred conventional additives include, for example, mannitol, a silane coupling agent, a water-soluble silicone and a polyelectrolyte.
- [0044] The electrolytic solution of the present invention can be prepared by mixing and dissolving the above-described various components in an arbitrary sequence according to a conventional procedure or a modified procedure thereof. For example, the electrolytic solution can be simply prepared by preparing a solvent containing a highly increased amount of water as a mixture of an organic solvent and water, and optionally dissolving an electrolyte, a nitro compound and optional additives in the resulting solvent.
- [0045] According to the present invention, there is also provided an electrolytic capacitor, preferably an electrolytic

capacitor comprising a capacitor element formed of an anode foil, a cathode foil opposed to the anode foil and a separator disposed between the anode foil and the cathode foil, and the electrolytic solution of the present invention.

[0046] The electrolytic capacitor of the present invention is more preferably an aluminum electrolytic capacitor, and most preferably an electrolytic capacitor comprising:

a capacitor element formed by winding an anode foil consisting of an aluminum foil and an anodized film appearing on the surface of the aluminum foil, and a cathode foil made of the aluminum foil, via a release paper, so that surfaces of both foils face each other;

an electrolytic solution of the present invention;

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a case or casing containing the capacitor element and the electrolytic solution; and an elastic sealant with which an opening portion of the case is sealed.

[0047] In the electrolytic capacitor of the present invention, because the electrolytic solution of the present invention is used, the functions of improving low-temperature stability based on a mixed solvent of an organic solvent and water, the hydrogen gas absorption function based on addition of a nitro compound, and prolongation of a working life and reduction of impedance based on inhibition of the hydration reaction due to use of a specific electrolyte, can be attained.

[0048] The electrolytic capacitor of the present invention is preferably formed in such a manner that a capacitor element is formed by winding an anode foil, wherein the surface of an etched aluminum foil is anodized, and a cathode foil made of the etched aluminum foil, via a release paper, so that surfaces of both foils face each other, and an electrolytic solution of the present invention are contained in a case, and an opening portion of the case containing the capacitor element is sealed with an elastic sealant.

[0049] Figure 1 is a sectional view showing one preferred embodiment of the electrolytic capacitor of the present invention, and Fig. 2 is a perspective view, enlarged partially in the thickness direction, which shows a capacitor element of the electrolytic capacitor shown in Fig. 1. Although the embodiment shown in the drawings is an electrolytic capacitor with a wound structure, various changes and modifications may be made in the electrolytic capacitor of the present invention without departing from the spirit and scope thereof. Of course, electrolytic capacitors other than the electrolytic capacitor with a wound structure are included in the scope of the present invention.

[0050] The illustrated electrolytic capacitor 10 is a chip-shaped aluminum electrolytic capacitor and has such a structure that a capacitor element 1 impregnated with an electrolytic solution is contained in a metal case 4 and an opening portion of the case 4 is sealed with a sealant 3. The capacitor element 1 contained in the metal case is in the form of a wound sheet-like laminate 20. The laminate 20 comprises, as shown in the drawing, an aluminum foil (anode) 21 having an oxide film 22 over the entire surface thereof, an aluminum foil (cathode) 23, a first separator (release paper) 24 interposed between these electrodes, and a second separator (release paper) 25. The first separator 24 and the second separator 25 may be the same or different, but are preferably the same in view of the production process and cost. The second separator 25 may be formed from a usual insulating film, if it is necessary. The capacitor element 1 is impregnated with an electrolytic solution.

[0051] In the illustrated electrolytic capacitor 10, the sealant 3 has a lead wire-penetrating hole for inserting a lead wire 2, thereby to conduct sealing, therein. The end of the opening portion of the case 4 is provided with a curl 14 to enhance a sealing strength of the sealant.

[0052] The electrolytic capacitor shown in Figs. 1 and 2 can be produced, for example, by the following procedure. First, an anode foil, wherein an oxide film is provided over the entire surface, by anodizing the surface, of a high-purity aluminum foil as a raw material, and a cathode film whose surface area is increased by etching the surface are made. Then, the resulting anode foil and cathode foil are disposed facing each other and a separator (release paper) is interposed between those films to form a laminate, thereby making an element with a structure obtained by winding this laminate, that is, a capacitor element. Subsequently, the resulting capacitor element is impregnated with an electrolytic solution and the capacitor element impregnated with the electrolytic solution is contained in a case (generally made of aluminum), as described above, and then an opening portion of the case is sealed with a sealant. Two lead wires are inserted into a lead wire-penetrating hole of the sealant, thereby to completely prevent leakage of the electrolytic solution.

[0053] The electrolytic capacitor of the present invention will be described in more detail hereinafter.

[0054] The aluminum foil used as the anode foil and cathode foil is preferably an aluminum foil having high purity of 99% or more. The anode foil can be preferably formed by electrochemically etching the aluminum foil, anodizing it to form an oxide film on the surface, and attaching a lead tab for connecting an electrode. The cathode film can be formed by etching the aluminum foil and attaching a lead tab for connecting an electrode. This cathode foil may not be anodized.

[0055] The capacitor element can be obtained by winding the anode and cathode foils, formed in the above steps, via the above-described release paper while the surfaces of both foils face each other.

[0056] The release paper used in the production of the capacitor element is not specifically limited, but is preferably

a paper produced by using as a raw material a naturally produced cellulose material, for example, Manila hemp and raw pulp. As the release paper, for example, there can be advantageously used a paper produced by passing the raw pulp through a dust removing process, a washing process, a beating process and paper-making process. A paper derived from synthetic fibers can also be used, however, such a paper is not preferred because it is inferior in heat resistance and corrosion of the capacitor is caused by halogen ions contained in the paper.

[0057] The sealant used in the electrolytic capacitor of the present invention can be formed from various materials used usually as far as the material has high hardness and proper rubber elasticity, and it is also impermeable to an electrolytic solution and has good airtightness for the sealant. Preferred sealant material includes, for example, elastic rubber such as natural rubber (NR), styrene-butadiene rubber (SBR), ethylene-propylene terpolymer (EPT), and isobutylene-isoprene rubber (IIR). The isobutylene-isoprene rubber (IIR) is preferably used because the airtightness is high and the electrolytic solution does not penetrate in the form of vapor. Vulcanized IIR having more excellent heat resistance, for example, sulfur-vulcanized, quinoid-vulcanized or resin-vulcanized IIR is used more preferably, and the resin-vulcanized IIR is particularly preferred.

[0058] In the practice of the present invention, a hybrid material obtained by laminating a resin material plate having sufficient airtightness and strength (e.g. fluorine-contained resin plate such as PTFE plate) can be advantageously used in place of the above-described sealant material.

Examples

[0059] The following Examples further illustrate the present invention in detail. Note that these examples are to be construed in all respects as illustrative and not restrictive.

Example 1

5 [0060] An aluminum electrolytic capacitor with a wound structure was produced in accordance with the following procedure.

[0061] First, an aluminum foil was electrochemically etched, followed by anodizing to form an oxide film over the entire surface of the aluminum foil, and then a lead tab for connecting an electrode was attached to make an aluminum anode electrode. Another aluminum foil was also electrochemically etched and a lead tab for connecting an electrode was attached to make an aluminum cathode electode. Subsequently, a capacitor element was made by interposing a separator (release paper) between the anode foil and the cathode foil, followed by winding. The capacitor element was impregnated with an electrolytic solution whose compositions are shown in Table 1 below and contained in an aluminum case with a base so that the lead tab for connecting an electrode protrudes out of the case, and then an opening of this case was sealed with an elastic sealant to make an electrolytic capacitor with a wound structure (10WV-1000 µF).

[0062] The specific resistance at 30°C of the electrolytic solution used in this example was measured to obtain measured values as described in Table 1 below. After an impedance at low temperature (-40°C) and an impedance at normal temperature (20°C) of the electrolytic capacitor thus obtained were measured, an impedance ratio (ratio Z) expressed as a ratio of the respective measured values was determined at different frequencies: 120 Hz and 100 kHz. As a result, measured values as described in Table 1 below were obtained. To evaluate characteristics of working life of the respective electrolytic capacitor, an initial value (characteristic value immediately after production of a capacitor) and a characteristic value after the capacitor was allowed to stand at high temperature (lapse of 1000 hours at 105°C) under application of a rated voltage were measured with respect to the capacitance, $\tan \delta$ and leakage current. As a result, measured values as described in Table 1 below were obtained.

45 <u>Examples 2 to 10</u>

[0063] The same procedure as in Example 1 was repeated, except that in this example, compositions of the electrolytic solution were changed as described in Table 1 below. The results of characteristic tests are summarized in Table 1 below.

Comparative Examples 1 to 3

[0064] The same procedure as in Example 1 was repeated, except that in this example, for the comparison purpose, a nitro compound was eliminated from the electrolytic solution and that compositions of the electrolytic solution were changed as described in Table 1 below. The results of characteristic tests are summarized in Table 1 below.

								_
5		•	at 105°C	Leakage Ocurrent	5		2.3	
			hours	tan 8			6.2	
10			After 3000 hours at 105°C	Capacitance tan [4]		•	869	•
15	•		90,	Leakage current [uA]			7.7	
			Initial value	tan A [8]			5.4	
20			Initi	Capacitance tan A [4]			1044	
25		Table 1	Ratio 2	100 kHz [-40/20°C]			4.6	
30			Rat	120 Hz [-40/20°C]			1.1	
35			Specific	resistance at 30°C [Ω·cm]			28	
40			positions of	electrolytic solution (% by weight)	Ethylene glycol 25.0 Water 68.0	um formate.	Hypophosphorous 0.4	n-nitroacetophenone 1.0
			Comic	elec (* }	Ethyle Water	Amac	Hypop acid	ր-ս

	Compositions of		Specific	Rat	Ratio 2	Initial	al value	e a	After 3000 hours at 105°C	hours	at 105°C
Example No.	Example electrolytic solution No. (% by weight)		resistance at 30°C (Ω·cm]	120 Hz [-40/20°C]	100 kHz [-40/20°C]	Capacitance tan A [4]		Leakage current	Capacitance tan Λ Leakage [μF] (%)	tan 8	Leakage current
	ne glycol	25.0									5
		68.0									
Example	im formate.	4.6									
,	Hypophosphorous acid	0.4	28	1.1	4.6	. 1044	5.4	7.7	869	6.2	2.3
	n-nitroacetophenone	1.0									
	Nitrobenzoic acid	1.0									
		20.0									
Person 10		60.0			•						
trambra -		16.4		,	,						
7	Sulfamic acid	1.6	97	7.1	4.4	1034	S. 4	7.2	906	6.2	2.3
	Nitrophenol	1.0									
	n-nitroacetophenone	1.0									
	ne glycol	15.0			-						
Example Water		60.0									
~		23.0	23	1.1	3.9	1025	5.3	7.0	902	-	0.0
,	Nitrobenzoic acid	1.0								<u> </u>	;
	Nitrophenol	0.4						,			
	ı	22.0							-		
Frame		50.0									
1	ate	25.0		•	c c	-	. 1			<u> </u>	,
4	Benzenesulfonic	-,	17	1.1	ב יי	1020	5.5	6.9	916	- 0.9	5.0
	acid	9.					-				
	Nitrophenol	2	•								

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5	•			
10				
15				
20				~
25				Continued
30				Table 1 (
35				
40				
45			•	

		Γ	Specific	Rat	Ratio 2	Initi	Initial value	90	After 3000 hours	hours	at 105°C
Example No.	Example Compositions of No. (% by weight)		resistance at 30°C [Ω·cm]	120 HE [-40/20°C]	kHz)/20°C]	Capacitance tan	tan b [8]	Leakage current [[A]	Capacitance (µF)	tan h [4]	Leakage current (µA)
Example Rehyle S Ammoni	ne glycol um borate	48.0 40.0 11.0	161	1.2	. 9'5	1024	9.7	6.2	932	9.5	1.9
	Nitrobanzolc acid	1.0									
		34.0									
Example		30.0			-	•					
1 2	ıte	14.6	29	1.0	3.7	1010	5.4	6.1	929	6.2	5.0
•	Phosphoric acid	4.0							•		
	n-ni troacetophenone	1.0									
	ne glycol	60.09			•						
		20.0	,								
Example		18.6	58	1.0	3.6	1003	6.2	6.3	933	7.0	2.1
۲	7 Boric acid	9.0	-								
	Nitrobenzoic acid	1.0									
		62.0									
		27.0									
Example	um adipata .	9.0	94	1.0	3.8	1005	7.1	6.2	940	7.9	2.1
&	Ni traace tophenone	1.0		•			_				
	Nitropenol	1.0									
	ne glycol	40.0									
0.000	Water	40.0	-	•	•	9	6	•			,
DT CONTRACT	Ammonium glutarate	19.0	<u> </u>	?	P.	1010	n 0			•	7.7
		1.0									•
	Ethylene glycol	50.0									
		39.4			•						
Example	Example Ammonium adipate	8.5	89	1.0	3.7	1013	6.4	6.4	942	7.0	2.3
01	Sulfamic acid	4.0									
	Nitrophenol	1.0				-					

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	105°	Appearance		of safety-	lution		of safety-	lution		of safety-	lution	
	After 3000 hours at 105°r	Leakage	(\	peration	y gas evo		peration	y gas evo		peration	Y gas avo.	
	0000	ten ()		98,	o pe	nr. 8	, s	0 P		o's	d De	B Z N
	After	Capacitance tan & Leakage Capacitance tan & Leakage durrent (if) current (if)		In all samples, operation of safety-	6.3 Vent was caused by gas evolution	WALET SOU DOUES	In all samples, operation of safety-	Vent was caused by gas evolution	אברוויט מכי שמודא	In all samples, operation of safety-	vent was caused by gas evolution	within 250 hours
	97	Leakage	3					7:0		. ,	6.9	
	Initial value	tan' 5 [8]			?						4.7	
inuedl	Initia	Capacitance [jiF]		0001	000		7617	5			1023	
Table 1 (continued)	Ratio 2	120 Hz 100 kHz Capas [-40/20°C] [-40/20°C] [HF]		1 75			7			•	, ,	
Table	Rat				;		1.1)			? .	
	Spacifia	resistance at 30°C (O.cm)		8.5			40			20	;	
		ution	0,00	30	10.0	45.0	40.0	15.0	30.0	30		3
	Compositions of	Example electrolytic solu (% by weight)	Comp. Ethylane glygol	Water	1 Ammonium adipate	Ethylene glycol	Water	2 Ammonium adipate	Ethylene glycol	Water	3 Ammonius adinate	0104777
		Example No.	Comp.	Example Water	-1	Comp.	Example Water	2	Comp.	Example Water	· м	

[0065] As is apparent from the results described in Table 1, the resistivity of the electrolytic solution except for Example 5 is almost the same as that of the Comparative Examples and the specific resistance is smaller than that of

a conventional electrolytic solution. Although the specific resistance of the electrolytic solution of Example 5 shows a large value such as 161 Ω • cm, the electrolytic capacitor is substantially comparable with a conventional electrolytic capacitor and is suited for practical use when generally judged considering other characteristics. Accordingly, the electrolytic capacitor made by using the electrolytic solution of the present invention can realize a lower impedance than a conventional electrolytic capacitor, or can realize a low impedance which is equivalent to that of a conventional one.

[0066] It has been found that the electrolytic capacitor using the electrolytic solution of the present invention has a small ratio Z and that the ratio Z at a high frequency such as 100 kHz is particularly reduced as compared with those of the Comparative Examples. This fact shows that the electrolytic capacitor using the electrolytic solution of the present invention exhibits good low-temperature stability over a wide frequency range.

[0067] Particularly, the electrolytic capacitor using the electrolytic solution of the present invention shows stable characteristics under application of a rated voltage even after it was allowed to stand at high temperature (lapse of 3000 hours at 105°C) by adding the nitro compound in the electrolytic solution in the amount ranging from 0.01 to 3% by weight, and the capacitor itself was not broken by gas generation. On the other hand, it became impossible to use all electrolytic capacitors of the Comparative Examples using the electrolytic solution containing no nitro compound because a safety-vent operated as a result of expansion of the case caused by hydrogen gas generation at an initial stage before a lapse of 3000 hours. This fact shows that a working life of the electrolytic capacitor can be easily prolonged according to the present invention.

Examples 11 to 19

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[0068] The same procedure as in Example 1 was repeated, except that in this example compositions of the electrolytic solution were changed as described in Table 2 below to confirm the effect of simultaneous addition of a chelate compound and a nitro compound. As is summarized in Table 2 below, satisfactory results could be obtained. In Table 2 below, the test results of Comparative Examples 1 to 3 are also described.

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				Y ATOBY	7 8						
	Compositions of	Specific		Ratio Z	Initi	Initial value	ne ne	After	1000	After 1000 hours at 105°C	105°C
Example No.	No. (% by weight)	resistance at 30°C [\Omega_cm]		120 Hz 100 kHz Capa [-40/20°C] [-40/20°C] [µF]	citance	tan d [%]	Leakage current	Capacitance tan h Leakage [[iE] [w] current	tan d	Leakage current	Leakage current Appearance
Comp.	Ethylene glycol 60.0	0.						To all comm			
Example		30.0	1.3	36.1	1008	10		and all samples, operation of	, ,	peration	
	1 Ammonium adipate 10	10.0					1	evolution within 500 hours	ithin	500 hours	2 2
Comp.		0.						In all samples operation of	lea.	peration	90
Example	Water 40.0	.0	1.1	6,6	1014	5.7	1.9	safety-yent the Caused by the		Total Pool a	
~	2 Ammonium adipate. 15.0	0.			•			evolution within 250 hours	Lthin	250 hours	
Comp.	Ethylana glycol 30.0	0.						In all camples operation of	9	Coration	
Example	Water 50.0	.0 20	1.0	7.9	1023	4.7	6	safety-vent was caused by mas		normand by	
-	3 Ammonium adipate 20.0	0.						evolution within 250 hours	Lehin	250 hours) 1
	Ethylene glycol 25.0	0.									
	Water 69.4	7.				_					
		4.0				_					•
Example 11		0.4 21	1.1	9.1	1044	5.5	7.8	919	S.8	2.5	88 C. 1.9 -
		8.0					•				raccory
		0.4									
	ne glycol 2	0.									
-	Water 59.2	.2									
Example	Example Amnonium 17 8										
12		92	1.1	4.4	1036	5.4	7.3	922	0.9	2.3	Sacis-
!		1.6			-			:			factory
	Nitrophanol 1.	1.0									
		0.4									_

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			Tabl	2 (0	Table 2 (continued)			
	l	Specific	Ratio 2		Initia	Initial value	After 1000 hou	1000 hou
strions or rolytic sol	utton	rolytic solution	120 Hz 100 kHz Capacitance ton N	2HX 00	Capacitance	Leaka	Leakage Capacitance ten X Le	N Le

		Specific	Rat	Ratio 2	Initi	Initial value	10	After	1000 h	After 1000 hours at 105°C	J. 501
Example	Example compositions of	resistance	120 Hz	100 kHz	100 kHz Capacitance tan & Leakage	tan Ö	Leakage	Capacitance tan 8 Leakage	tan A	Leskage	
Š	(% by weight)	[D.G]	[-40/20°C]	[-40/20°C] (-40/20°C] [[HE]	[311]	<u>×</u>	current [IA]	[h]	3.	current [IIA]	Appearance
	Ethylene glycol 15.0										
											_
Ardwox.	Andrea Ammontum adipate 24.4	23	1.1	3.9	1028	5.3	7.1	925	5.9	2.5	95t18-
;	T										factory
											
	Ethylene glycol 24.2										
	Water 50.0										
9										_	
C x camp 4 c	Sampia Benzeneguifonic	21	1.1	3.8	1021	5.2	6.9	930	8.8	7.7	satis-
``	ectd							•			ractory
	p-nitrobenzoic acid 0.8										
	DTPA 0.4										-
	Ethylene glycol 55.0									ĺ	
	Water 28.0	•	•								-
Example	Example Ammonium sulfamate 14.0		•		-	•	,		•		satis-
1.5	Phosphoric sold 2.0	ŝ	1.0	7.7	1003	4.0	2.9	926	0.9	2.5	factory
	Nitrophenol 0.6										• •
	EDTA 0.4										-

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Table 2 (continued)

		Specific	Rat	Ratio Z	Initia	Initial value	9	AFFOR	1000	After 1000 bones of 10500	0.30
Example No.	Example compositions or No. (% by waight)	resistance at 30°C	120 Hz [-40/20°C]	100 kHz (-40/20°C)	Capaci	tan N	Leakage	Capaci	tan N	Leakage	Appearance
	C 03 . [20::[= 000[:0446	(mo . 24)					(A)		2	[N]	
	a yayear										•
_				_							
Example	_		,	,							444
7.9	16 Boric acid 0.4	25	1.0	۵. ۳	1002	6.1	6.4	944	6.7	2.4	80118
			-			_					ractory.
	Ethylene glycol 62.0										
					•					-	
Example		-					•				
11	17 EDTPO 0.1	92	0.1	8. 8.	1003	7.0	6.5	942	7.6	2.4	SECTOR S
	Nitrobanzoic acid 1.2										I accord
•	EDTA 0.4										
	ne glycol										
Trend	Water 40.0										
Brdingra	Ammonium glutarate 19.6	39	1.0	3.6	1018	8,5	6	637	v		satis-
; 	Nitrophanol 1.2						!		;	:	factory
	EDTPO . 0.4									-	
	Ethylene glycol 48.8								Ì		
	Water 40.0							_			
Example	Example Ammonium adipate 9.2		,								1 6 6
19		20	0.1	3.7	1014	4.9	6.2	943	7.0	2.5	factory.
					-						_

Examples 20 to 29

[0069] The same procedure as in Example 1 was repeated, except that in this example compositions of the electrolytic solution were changed as described in Table 3 below to confirm the effect of simultaneous addition of saccharides and a nitro compound. As is summarized in Table 3 below, satisfactory results could be obtained. In Table 3 below, the test results of Comparative Examples 1 to 3 are also described.

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					7724							-
	Composit Cons of	•	Specific	Rat	Ratio Z	Initia	Initial value	97	After	1000 h	After 1000 hours at	105°C
Example No.	electrolytic solution (% by weight)	ton	resistanda at 30°C [N·cm]	120 Hz [-40/20°C]	120 Hz 100 kHz Capa [-40/20°C] [-40/20°C] [[IF]	cítance	tan 8	Leakage current	Capaci [µF]	tan b %	Leakage Current	Appearance
Comp.	Ethylene glycol	60.0							In all samples copration of andotter		in the second	2000
Example	Water	30.0	88	1.3	36.1	1008	7.0	5.5	Vent ves caused by	ייייייייייייייייייייייייייייייייייייי	das avolution	Intion
-	Ammontum adipate	10.0					-		within 500 hours	lours 2		
Comp.	Ethylene glycol	45.0							In all samples coeration of estatu-	0	or ar i	of oafohir-
Example		40.0	40	1.1	9.7	1014	5.7	6.1	vent was caused by cas avolution	y y besi	CAB BAD	lution
7	2 Ammonium adipate	15.0							within 250 hours	OUEB		
Comp.		30.0							In all samples constation of safety-	90	Grafion	of parotu-
Ехамріе		50.0	20	1.0	7.9	1023	4.7	6	Vent was caused by one evolution	yd bee	000000000000000000000000000000000000000	Tables 10
n		20.0							within 250 hours	Ours		
	e glycol	23.2										I
		70.0									•	_
		4.4							•			
o Tdwaxa	Galactose	1.0	22	1,1	4.6	1043	. S	7.7	918	0	2.5	satis-
?	Nitrophanol	1.0									}	factory
	Hypophosphorous	0		·			_	•				
	Ethylene glycol	5.					Ī					
		60.0	-	-								
Ежамріа	Lan Pre-	17.8				200	-					satis-
51	Fluctose	1.0		:	:		7	*:	176		?;	factory
	Nitrobenzoia acid	0.8										
	- 1	1.0										
	ne glycol	14.2					Ī					
		60.0									•	
Example	ţ,	23.B	2.4	-	0	1001			-			satis-
72	Dinitrobenzoic	1.0	•	•	•		·	?	776	J	7.7	factory
	release release	-									•	

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		Specific	Rat	Ratio 2	Initi	Initial value	87	After	1000 h	After 1000 hours at 105°C	105°C
Example No.	Example compositions or alectrolytic solution (% by weight)	resistance at 30°C	120 Hz [-40/20°C]	100 kHz [-40/20°C]	Capacitance tan & Leakage	tan ö	Leakage current	Capacitance tan 0 Leakage	tan d	Leakage current	Appearance
		(S2 · Ga)					(jd)	, p	/e	[HA]	
	Ethylane glycol 20.8	89.									
	Water 50.0	0.									
	Ammonium succinate 24.8	8.									•
4	Glucose	1.0 22	1.1	3.8	1020	5.3	8.9	930	9.3	2.2	satis-
3.		3.0									factory
•		. 4	-		•						
	acid					•					
	Ethylene glycol 48.7	6.									
	Water 39.8	80	_								
erdwerd.		9,0 162	1.2	5.6	1014	8.6	6.2	933	9.4	2.1	satis-
	-	1.5					•	-			factory
		1.0			;						
	ie glycol	.2	•								
	Water 30.0	0									
Example	Ammonium sulfamate 13.6	89					•		,		satis-
25	Fluctose 1.	1.0	7	· ·	8001	υ υ		756		2.2	factory
	Nitrobenzoic acid 1.	1.5		•				-			•
	Phosphoric acid 0.5	5			٠			-			
	Ethylana glycol 59.2	2									
	Water . 20.0	0									
o Cuma A	Ammonium adipate 17.8	89									
26	of Glucose 1.0	65 0	1.0	3.6	1001	6.2	6.3	944	9	2.4	BAC18-
 :	Dinitrobenzoic 1 0	-									ractory
	world	•									
	Boric acid 1.0										

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Table 3 (continued)

		Specific	Ratio Z	2 0	Initia	Initial value	9	After	1000	After 1000 hours at 105°C	0.50
Example No.	Example Compositions of No. (% by weight)	resistance at 30°C [Ω·cm]	120 Hz [-40/20°C]	120 Hz 100 kHz Capa (-40/20°C) [µF]	7	tan 6 [%]	Leakage current	Capaci [µF]	tan 8 [%]	Leakage current	Appearance
Example 27	Example Water 28.0 Smonium adipate 9.3 Nitrobenzoic acid 0.8 Fluctose 1.0		1.0	я. В.	1002	7.0	7.0 6.2	944	7.6	2.4	satis- factory
Example 28	Exhylena glycol 38.2 Water 40.0 28 Ammonium glutarate 18.8 Nitroacetophenone 2.0 Galactoso 1.0	40	1.0	3.6	1018	N. es	6.3	937	6.4	2.1	satis- factory
Example 29	Ethylene glycol 47.7 Water Example Ammonium adipate 9.0 29 Sulfamic acid 0.4 Xulreobenzoic acid 2.5	69	1.0	r.e	1013	δ.	6.2	942	7.0	2.2	satis- factory

Examples 30 to 39

[0070] The same procedure as in Example 1 was repeated, except that in this example compositions of the electrolytic solution were changed as described in Table 4 below to confirm the effect of simultaneous addition of hydroxybenzyl alcohol, glutamic-diacetic acid and the like as well as a nitro compound. As is summarized in Table 4 below, satisfactory results could be obtained. In Table 4 below, the test results of Comparative Examples 1 to 3 are also described.

Table 4

	Compositions of		Rat	Ratio 2	Initi	Initial value	ne ne	After	1000 h	After 1000 hours at 105°C	105°C
Example No.	No. (% by weight)	on at 30°C	120 Hz [-40/20°C]	120 Hz 100 kHz Caps [-40/20°C] [-40/20°C] [µF]	oitance	tan 6 %		Capacitance tan 5 Leakage (IRE) (ILE)	ce tan 8 %	Leakage current	Leakage Current Appearance
Comp.	Comp. Ethylene glycol 6	0.0					T	In all samples operation of eaforts.	8	aration,	- Section 1
Example Water		30.0	E	36.1	1008	2.0	6.5	Vent was caused by as evolution			int (an
-	Ammonium adipate 10.	0.0						within 500 hours	ours		
Comp.	Ethylene glycol 4	5.0						In all samples operation of sefetu-	6	pration	f anfotus
Example Water	Water	0.0	1.1	9.7	1014	5.7	6.1	Vent was caused by das evolution	and bea	CAS BAD	urion .
. 2	Ammonium adipate 1:	5.0						within 250 hours	ours		
Comp.	Ethylene glycol	0.0						In All samples operation of osfetue	8	eration o	f on fature
Example Water		50.0	1.0	7.9	1023	4.7	6.9	Vent was caused by das avolution	אל הפני	Cas ago	ition a
	Ammonium adipate 20	0.0			•			within 250 hours	ours		
	Ethylene glycol 2	24.0									
		0.6									
	Ammonium formate	4.6		_							
Example	Example Hypophosphorous	3.4	-	4					1	,	satis-
9	Dinitrobancoic		1	,	7	v	:	A 1 A		2.5	factory
	acid	1.2		•				•			
	Hydroxybenzyl	0.0						-			

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<i>30</i>	
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Table 4 (continued)

			-									
		Specific	707	RAC.	Ratio Z	IUICI	INICIAL VALUE	40	Arter .	1000	Arter 1000 hours at 105 C	. U.S. C
Example No.	Example Compositions of state No. (4 by weight)	on at 30°C	801	120 Hz [-40/20°C]	120 Hz 100 KHZ Capa [-40/20°C] [-40/20°C] [µF]	Capacitance (µF)	tan d %	Leakage current (IA)	100 kHz Capacitance tan 0 current [iP]	tan b %	Leakage current [IA]	Appearance
	Ethylene glycol Water	17.7						·				
скамруе		16.8		,				,				satis-
31	31 Sulfamic acid	1.6			٠ ٠	1034	υ 4	7.2	920	0.9	۲. ۲. ع	factory
	Nitrophenol	2.5										
	Glutamic-diacetic acid	1.4										
	ene glycol	14.2	<u> </u>									
		0.09	_									
xample		23.4		-		10.5		,	623	0	,	satis-
32				:	•	222	ì.		3	;	•	factory
	Glutamic-diacetic acid	1.4										
	ene glycol	20.8	T									
		50.0				•						
		-									_	
	•	24.8								:		4
PTdurx	Benzosulfonic	22		1.1	3.8	1020	5.3	8.9	930	5.9	2.2	88418
3	acid	, —				•						Lactory
	Nitrobanzoic acid	2.0										
	Mydrobensyl	2.0	-						•			

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		٠			able 4 (Table 4 (continued)		·					
-	Compositions of		Specific	Ra	Ratio Z	Initi	Initial value	91	After	1000 h	After 1000 hours at 105°C	.05°C	
Example No.	Ryampie electrolytic solution No. (1 by weight)		resistance at 30°C [$\Omega \cdot cm$]	120 Hz [-40/20°C]	100 kHz [-40/20°C]	Capacitance tan [µF] [%]	tan h [%]	Leakage current (IA)	Capacitance (µF)	tan 1) [%]	Leskage current	Appearance	
	Ethylene glycol Water	44.7	•										
0 (0 0 0 0	Ammonium boxate	9.8	. •						•				
34	Dinitrobanzoic acid	1.5	191	1.2	10 10	1024	8.7	6.2	942	9.3	2.1	satis- factory	
	Hydroxybanzyl alcohol	0.	•		·.								
	e glycol	52.2											
	Annonium	2 .							•				
Example	Example sulfanate	. ·	ñ	1.0	3.7	1010	. 47	6.1	939	1.	2	satis-	
3 	Nitroscetophenone	, 0		•								factory	
	2	2.6				•							
-	Ethylene glycol	57.2		·						:			
		20.0											
•	ipate	15.0											
Example	Example Mitrophenol	3.0	63	1.0	9.6	1003	6	4	776	. u	4	satis-	
۹ 	Hydroxybenzyl alcohol	2.6	-	:		}	;	?	•	;	•	factory	
	Glutamic-diacetic	,										_	

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10	•			
15				
20				
25	•		•	continued)
30				Table 4 (continued)
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		Specific	RACI	Ratio 2	Initia	Initial value	97	After	1000 h	After 1000 hours at 105°C	105°C
Example No.	Example Compositions of alectrolytic solution (% by weight)		120 Hz [-40/20°C]	100 kHz Capa [-40/20°C] [µF]	citance	tan 8 %	Leakage current [µA]	Capac (IE)	tan 8 [%]	Leakage current [IA]	Leakage current Appearance [IA]
	Ethylene glycol 59.2										
	Water 27.0										
	Ammonium adipate 9.3										
Example	Example Dinitrobenzoic 1.5		•	r		,	,				satis-
37	BOLD	n M	> -	B.	· cony	:	y .	٠ ٢	``	F. 7	factory
	dlutarate 1.5										-
	Glutamic-diacetic 1.5									-	
	Ethylene glycol 39.6	-									
	Water 40.0										
Example	Ammonium 19.0	96	1.0	9,6	1018	εΛ 60	e .	. 937	4		satis-
음 -	Nitrophenol 0.4								;	!	factory
	Hydroxybenzyl 1.0										
	Ethylene glycol 48.7										
•	Water 39.4							-			
÷	Ammonium adipate 9.0										
Example	Sulfamic acid 0.4		,			,	•		•	•	satis-
39		B 0	>. ₹	ì	1013		٠. ه	2 96 2	0.	2.3	factory
			,					-		-	
	Hydroxybenzyl 1.0			_							٠

Examples 40 to 49

[0071] The same procedure as in Example 1 was repeated, except that in this example compositions of the electrolytic solution were changed as described in Table 5 below to confirm the effect of simultaneous addition of a nitro compound and gluconic lactone. As is summarized in Table 5 below, satisfactory results could be obtained. In Table 5 below, the test results of Comparative Examples 1 to 3 are also described.

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		Specific	Rati	Ratio Z	Initia	Initial value	10	After	3000 h	After 3000 hours at 105°C	105°C
Example	Example compositions of resistance No. No. No. No. 130°C	(% at 30°C		100 kHz	Capacitance	tan û	Leakage	Capacitance tan & Leakage Capacitance tan & Leakage	tan 8	Leakage	900
	by weight)	[Q·cm]	[-40/20°C]	(-40/20 C] (-40/20 C) (µF)	(hF)	3	[hh]	[HE]	Ξ	(A)	
Comp.	Ethylana glycol	0.09						In all samples, operation of safety-	39, ope	Bration	of safety-
Example		30.0	1,3	36.1	1008	7.0	6.5	6.5 vent was caused by gas evolution	sed by	gas evol	lution
-1	0	10.0						within 500 hours	ours		
Comp.	Ethylene glycol	45.0						In all samples, operation of safety-	18, ope	aration c	f safety-
Example		40.0	1.1	9.1	1014	5.7	6.1	vent was caused by gas evolution	red by	gas evol	ution
2	Ammonium adipate	15.0						within 250 hours	urs		
Comp.	Ethylene glycol	30.0						In all samples, operation of safety-	st, ope	sration o	£ safety-
Example		50.0	1.0	7.9	1023	4.7	6.9	vent was caused by gas evolution	led by	gas evol	utton
3		20.0						within 250 hours	ours	ı	
	e glycol	25.0									
		0.89			•						
	Ammonium formate	1.4									
Example 40	Example Hypophosphorous	0.4	1.1	9.4	1044	5.6	7.7	868	6.4	2.3	satis-
	•	0.2				_					•
	n-nitroacetophenone	1.0									
_	n-nitrobenzoic acid). 				•			_		•

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Table 5 (continued)

	Compositions of	Specific	٥	Rat	Ratio 2	Initia	Initial value	en	After	3000 h	After 3000 hours at 105°C	105°C
Example No.	No. (1 by weight)	resistance at 30°C (Ω·cm)	90	120 Hz [-40/20°C]	120 Hz 100 kHz Capa	Capacitance tan & Leakage Capacitance tan & Leakage (18) (18) Current Appearance	tan 8 (8)	Leakage	Capacitance (HF)	tan ñ	Leakage current	Appearance
	Ethylene glycol	20.0	-					5			(<u>4</u>	
	Water	0.09										
Feen	Ammonium glutarate	16.2										
T T	Al Sulfanio soid	1.6	_	1.1	7	1034	ď	,	o c	,		satis-
;	Gluconic lactone	0.2		1	;		1	y.,	006	?	2.3	factory
	Nitrophenol	1.0					_					•
	n-nitroacetophenopne	1.0					_				_	_
	Ethylene glycol	15.0	-				T					
	Water	60.09			٠.							•
Example	Example Ammonium adipate									_	-	
\$		0.2			g. 6	1025	5.4	7.0	902	6.2	2.0	satis
		1.0		_								ractory
	Natrophenol	1.0		_					•	<u></u>		
	ne glycol	22.0	-				T				1	
		20.0			_		_					
Example	Example Ammonium succinate											•
£	acid	0.4 21	_		3.8	1020	5.5	9. ₉	916	6.0	2.0	satia-
	Gluconic lactone	0.5	_	-					•			ractory
	Nitrophenol	2.6			_							

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Table
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	-	Specific	P. Br	Ratio 2	Initi	Initial value	9	After	3000 h	After 3000 hours at 105 C	105°C
Example	Example Compositions of stample sleetrolytic solution (*)			100 kHz	100 kHz Capacitance tan 6	tan ń	Leakage	Leakage Capacitance tan A Courset annearance	tan ù	Leakage	Acceptance
2	No. by weight)		[-40/20°C]	-40/20°C] [-40/20°C] [[IE]		Ξ	(VI)	[.htg.]	Ξ	[hA]	
	Ethylene glycol 48	48.0									
		40.0									1 4 6
exambre	Example Ammonium borate . 10	10.8 161	1.2	5.6	1024	8.8	6.2	932	9.6	1.9	factory
;		0.2			•						
	Nitrobenzoio soid 1	1.0									
		54.0									
		30.0									
Example	Example Ammonium sulfamate 14	9.	,	,	0.01	4		0.0	4	•	satis-
45	Phosphoric acid 0	0.4	?	; ;	2		•		;	<u>.</u>	factory
	Gluconic lactone 0		_							٠	
	n-nitroscetophenone 1	1.0									
		0.09								-	•
	Water 20	20.0									
Example	Example Ammonium adipate 16	16.4	-	7.	.001	,	~	ננס	-	1.6	satis-
46		0.4	3) 1			;	7	?	:	factory
	Gluconic lactone 0	7	•								
	Nitrobantoic acid	0				_			_		

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Compos. (% by (% by Ethyles Water Ammonis Clucon n-nitz Mitropi Ethyles Mitropi Mitropi Mitropi Mitropi Mitropi Ammonis Gluconi Gluconi Gluconi Gluconi Gluconi Gluconi Gluconi	1 1 2 2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3		8 8 S S	120 Hz [-40/20°C] 1.0 1.0 1.0	3.6 3.6 3.7	Initia Capacitance (µF) 1005 1018	1 valu (4) (6.9 6.5	Leakage current [[M] 6.2 6.3 6.3	After 3000 t (µF) 940 937	1 (8) 7.7 7.7 7.1	Leakage current [lux]	s (continuea)	Specific Ratio 2 Initial value After 3000 hours at 105°C	tion restatance 120 w. 100 kHz Capacitance han N Leakage Capacitance han N	at 30°C . 140 AE	[[-40/20°C1 [-40/20°C1 [[[[]]]]]]	[CO-CM] [-10/20 C] [-10/20 C] [10/2] [10/2] [10/2]	Ethylane glycol 62.0	Mater 27.0	un adipata . 6.8	0.2 35 1.0 5.8 1005 6.9 6.2 540 7.1	n-nitroacetophanone 1.0	Ethylena glycol 40.0	40.0	18.8 40 1.0 3.6 1018 5.8 6.3 937 6.4	0.5	Nitrobensoic acid 1.0	Ethylene glycol 50.0	Water 39.4	0.0	7.7	Gluconic lactons 0.2	
Specific Ratio 2 Initial value After 3000 hours teststance at 30°C [-40/20°C] [-40/20°C] [F] [F	Specific Ratio 2 Initial value Affer 3000 hours resistance at 30°C [-40/20°C]	Table 3 (Colltinued) After 3000 hours Ratio 2 Initial value After 3000 hours 120 Hz 100 kHz Capacitance tan \$\hat{1.0}\$ Capacitance Capacitanc	Initial value After 3000 hours citance tan % Current [µE] [µE] [µE] [µE] [µE] [µE] [µE] [µE]	Initial value After 3000 hours citance tan % Current [µE] [µE] [µE] [µE] [µE] [µE] [µE] [µE]	Initial value After 3000 hours citance tan % Current [µE] [µE] [µE] [µE] [µE] [µE] [µE] [µE]	value After 3000 hours at 105°C tan % Current [45]	Leakage Capacitance tan Leakage Capacitance tan Capa	After 3000 hours at 105°C Capacitance tan h Current [48] [48] [14] [14] 940 7.7 2.1 937 6.4 2.2 937 5.4 2.3	18) (IM) 7.7 2.1 7.7 2.2 6.4 2.2	Leakage current [liA]					Appearance					setis-	factory			4	factor.	Taccord				satis-	factory		

Examples 50 to 59

[0072] The same procedure as in Example 1 was repeated, except that in this example compositions of the electrolytic solution were changed as described in Table 6 below to confirm the effect obtained by an arbitrary combination of various additives. As is summarized in Table 6 below, satisfactory results could be obtained. In Table 6 below, the test results of Comparative Examples 1 to 3 are also described.

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					Table	9			•			
	Compositions of		Specific		Ratio Z	Initia	Initial value	97	After	3000	3000 hours at 105°C	105°C
No.	No. (* by weight)		resistance at 30°C [Ω·cm]	120 Hz [-40/20°C]	100 kHz [-40/20°C]	Capacitance ten 3 Leakage Capacitance ten 3 Leakage Current (11F)	tan 3	Leakage	Capacitance (µF)	Ean (Leakage	Appearance
Comp.	Ethylene glycol	8						3			(IN)	
Example Water		30.0	88	1.3	ו אר	800	-		In all samples, operation of safety-	0 '607	peration	of safety-
	.um adipate	10.0	!	:	;	201	:	o o	Vent was caused by gas evolution	used b	y gas evo	lution
Comp.		45.0							200	a inou		
Example		0.0	•	1.1	7.6	101	,		in all samples, operation of safety-	163	peration	of safety-
7		15.0		:	:		;		vent was caused by gas evolution within 250 hours	1984 D	у дав вvо	lution
Comp.		30.0							מריוויון לים	100		
Example		50.0	20	1.0	7.9	1001	,	4	in all samples, operation of safety-		peration	of safety-
-		20.0				}	-		went was caused by gas evolution	0 0000	y gas evo	Lution
		24.0							257	500		
_		68.0										
		4										
	Mypophosphorous acid	٠.										
2	EDTA	0.5	ć	. '				•				
20	Glutamic-diacetic	-	87	1:1	9.	1044		7.7	868	6.1	2.3	98718
	aoid ,	·			•							ractory
	Gluconia lactone	0.2										
	n-nitroscetophenone	1.0										
	Mitrobenzolo acid	1.0										
	ne glycol	18.0										
		0.09										
	nrato	16.2			-							
Example	Sulfamic acid	1.6										
		0.1	26	1.1	4.4	1034	5.2	7.2	000	4	ŗ	satis-
	Fluctosa	1.0					!	<u>.</u>	2	?	7	factory
	Gluconic lactone	0.2										
	Nitrophenol	1.0		-				_			_	
	n-nitroacatophenone	1.0										_

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	•		
		•	
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-		Specific	Ratio	io z	Initial	l value	ue	After	3000	After 3000 hours at	105°C
Example No.	Example electrolytic solution No. / A L / L	resistance at 30°C	120 Hz	100 kHz	Capacitance tan S current	tan S	Leakage current	Capacitance tan 3	tan à	Laskage	Leakage current Appearance
	(* by waight)	(Q·cm)	[-40/20°C]	[-40/20°C]	(µF)	3	[H)	[µF]	3	[hh]	
	ana glycol	0.									
		s.				•					
	Ammonium adipate 22	22.8						-			
2000		2.0							•		
52		0.5	1.1	3.9	1025	5.5	7.0	902	6.3	2.0	satis. factory
	Gluconic lactona 0	0.5									
	_	1:0				-					
	Nitrophenol	1.0									
	7001 2	9.									
		0.		•							
	Ammontum	-						٠			
	inate	•					-	•			
	EDDA	1.0									•
S3	Glutamic-discetic	0.2	1.1	9.0	1020	5.4	B.9	918	6.2	2.0	gatis-
	BOLD BOLD BOLD FOR IT										
		4.0									
	Gluconia lactone 0.	0.2						-			
.		2.6								-	
	Ethylene glycol 46.7	۲.									
		0						-			
	Ammonium borate 10.8	69	•								-
Example EDDA		0.8	•				,			•	satis-
ş	benzyl	5.0	•	•	• • • • • • • • • • • • • • • • • • • •	h	· ·	336	'n		factory
		<u> </u>									
		0.5									
	Nitrobengoic acid 1.	1.0									

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able 6 (continued)

	Compositions of	Specific		Ratio Z	Initi	Initial value		After	3000	After 3000 hours at 105°C	105°C
Example No.	No. (1 by weight)	resistance at 30°C (Ω·cm)	120 Hz [-40/20°C]	ٺ	citance		Leskage current	Capaci (µF)	tan 8	Leakage	Appearance
-	ne glycol	52.0					3		T	5	
		0.0									
	Ammonium sulfamate 1	14.4									-
	Xylose	0.5			<u>ک</u>						-
Example EDTPO			,		7						
55	Glutamic-diacetic	0.5	0.1	3.7	1010	8.5	6.1	929	6.0	2.0	factory
	3178	_		- ,					_		-
		. 4									
	Gluconic lactone	0.5									
	none	1.0									
	ĺ	57.5									
		0.0								_	
	Ammonium adipate 1	19.4		_			-				
		1.5									
Example.	Example Hydroxybenzyl			-							
3,4		28	1.0	3.6	1003	6.1	6.3	613	6 9	,	satis-
;	atic				!		:	}	;	:	factory
	acid			_				•		-	
		0:4									
	Gluconic lactons	0.2	•		-		-				
	Nitrobenzoia acid						_				

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	•		Tab	Table ([continued]	Truned)						
	-	0.000	Ratio 2	2 0	Initia	Initial value		After	3000 h		105°C
xample No.	xample compositions of albeiton No. (4 by seight)	resistance at	120 Hz	100 KHZ	Capacitance tan A current	tan a	Laakage current	Capacitance tan (Leakago	ten is	Laskago	Leakago current Appearance
				1-40/40 67	/ lie /	2	[M]	llie J	٤	[W]	
	Ethylene glycol 57.5	٠,									
	Water 27.	0.									
	Ammonium adipate . 8.	8.8									_
		0.		•							
xample	xample Glucose 1.	1.0		ď	1006	ď	,	. 676			satis-
21	Hydroxybenzyl 6.	6.5	?	7	5	;		ŝ	?	:	factory
	c lactone	0.2									
	910	1.0									
		0.									-
ŀ	100/	6									
		.0									
	EDDA 1.	0									
	Fluctose 0.5	87						-			4
58 58	58 Glutamic-diacotic 0.6	40	1.0	3.6	1018	4.0	6.4	937	4.	2.5	factory
											•
	_	5	-								
	Nitrobanzolo acid 1.0	7 0		•							
	}	3									
	Water 39.4	4									
	Ammonium adipate 9.	0									-
	benzyl					•					
xample	xample alcohol										satis-
59	EDTPO 1.0	89 0	1.0	3.7	1013	9 9	•	942	7.4	2.3	factory
	Fluctose 0.5									,	
	Sulfamic acid 0.4										
	one	2								-	
		-		-							

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Comparative Examples 4 to 6 and Examples 60 to 62

[0073] The same procedure as in Example 1 was repeated, except that in this example the measurement of the characteristic value under high temperature conditions (application of rated voltage, lapse of 1000 hours at 105°C) employed in Example 1 was conducted under conditions (lapse of 6000 hours at 105°C) to confirm a further improvement in characteristics of working life. The results as described in Table 7 below were obtained.

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				Table 7	e 7		:				
	Company 1 tops of	Specific	RAE	Ratio 2	Initia	Initial value	92	After	0009	After 6000 hours at 105°C	105°C
Example No.	Example composition of a long solution No. (1 by weight)	resistance at 30°C (Ω·cm]	120 Hz [-40/20°C]	100 kHE CBPA [-40/20°C] [IIF]	Capacitance tan h Leakage [IF]	tan b [8]	Leakage ourrent [µA]	Capacitance (µF)	tan 3 [8]	Leakage current [IA]	tan 3 Leakage Appearance [%] [IIA]
Comp.		0.09						In all samples, operation of safety-	9305	eration	of safety-
Example		30.0	1.3	36.1	1008	7.0	6.5	vent was caused by gas evolution	sed by	gas eve	Jutton
•		10.0			•		•	within 500 hours	ours	•	
Comp.		45.0						In all samples, operation of safety-	98. 00	eration	of safety-
Example		40.0	1.1	9.1	1014	5.7	6.1	vent was caused by gas evolution	sed by	das evo	lution
<u>.</u>	5 Ammonium adipate 15	15.0						Within 250 hours	ours		
Comp.		30.0						In all samples, operation of safety-	98. 00	eration	of safety-
Example		50.0	1.0	7.9	1023	4.7	6.9	vent was caused by das evolution	ed by	GA3 8V0	lution
9		20.0						within 250 hours	Dura		
•		25.0									
		68.0									
Example	Example Ammonium formate 4	4.6	•	,	,,,,,				,	,	satis
8	ų		:	0	***	, ,	<u> </u>	ກ ກ	و و	2.1	factory
	9	0.0									
	[0.									
-	Water 60.0	0.									
Exampra C.	Ammontum adipate 23.0	.0 23	1.1	9.0	1025	5.3	7.0	699	8.2	9 1	satis-
3		1.0		•	٠.						factory
		1.0		_							
	Ethylene glycol 40.0	0.							T		
9		0.	•		9	v		-	•		satis-
62	Ammonium glutarate 19.0		?	9.	8 707	D.		250	7. 6	7:1	factory
أب	Nitrobenzoic acid 1	1.0				•					

[0074] In Table 7, Comparative Examples 4 to 6 respectively correspond to Comparative Examples 1 to 3, while Examples 60 to 62 respectively correspond to Examples 1, 3 and 9. As is apparent from the results, it becomes impossible to use all capacitors of Comparative Examples 4 to 6 using an electrolytic solution having added thereto no nitro

compound, whereas, capacitors of Examples 60 to 62 could be used even after 6000 hours had passed although a reduction in capacitance was recognized. Surprisingly, it has been found that characteristics of working life of the electrolytic capacitor are further improved by using a carboxylic acid or a salt thereof as an organic electrolyte in combination with an inorganic acid as an inorganic electrolyte.

Industrial Applicability

As described above, according to the present invention, there is provided an electrolytic solution, for use in an electrolytic capacitor, which has a low impedance and excellent low-temperature stability expressed by an a ratio of an impedance at low temperature to that at normal temperature, along with good characteristics of working life, and also it can exhibit an excellent hydrogen gas absorption function when an electrolytic solution contains a highly increased amount of water in its mixed solvent or when an electrolytic capacitor is used under high temperature conditions. According to the present invention, there is also provided an electrolytic capacitor with high reliability, which has a low impedance and excellent low-temperature stability, along with good characteristics of working life, and also it is free from defects due to presence of water used as a solvent, specially an aluminum electrolytic capacitor.

Claims

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- An electrolytic solution for use in an electrolytic capacitor, comprising a solution containing a solvent consisting of 20 to 80% by weight of an organic solvent and 80 to 20% by weight of water, and at least one electrolyte selected 20 from the group consisting of a carboxylic acid or a salt thereof and an inorganic acid or a salt thereof, having added thereto at least one nitro compound selected from the group consisting of nitrophenol, nitrobenzoic acid, dinitrobenzoic acid, nitroacetophenone and nitroanisole.
- The electrolytic solution for use in an electrolytic capacitor according to claim 1, wherein the nitro compound is a 25 combination of two or more nitro compounds.
 - The electrolytic solution for use in an electrolytic capacitor according to claim 1 or 2, wherein the nitro compound is added in the amount of 0.01 to 5% by weight based on the total amount of the electrolytic solution.
 - The electrolytic solution for use in an electrolytic capacitor according to any one of claims 1 to 3, wherein the organic solvent is a protic solvent, an aprotic solvent, or a mixture thereof.
- 5. The electrolytic solution for use in an electrolytic capacitor according to any one of claims 1 to 4, wherein the carboxylic acid or salt thereof is selected from the group consisting of formic acid, acetic acid, propionic acid, butyric 35 acid, p-nitrobenzoic acid, salicylic acid, benzoic acid, oxalic acid, malonic acid, succinic acid, glutaric acid, adipic acid, fumaric acid, maleic acid, phthalic acid, azelaic acid, citric acid and hydroxybutyric acid, as well as ammonium, sodium, potassium, amine and alkyl ammonium salts thereof.
- 6. The electrolytic solution for use in an electrolytic capacitor according to any one of claims 1 to 4, wherein the inor-40 ganic acid or salt thereof is selected from the group consisting of phosphoric acid, phosophorous acid, hypophosphorous acid, boric acid, sulfamic acid, as well as ammonium, sodium, potassium, amine and alkyl ammonium salts thereof.
- 7. The electrolytic solution for use in an electrolytic capacitor according to any one of claims 1 to 6, further comprising 45 at least one additive selected from the group consisting of the following group:
 - (1) a chelate compound,
 - (2) saccharides.
 - (3) hydroxybenzyl alcohol and/or L-glutamic-diacetic acid or a salt thereof, and
 - (4) gluconic acid and/or gluconic lactone.
 - 8. An electrolytic capacitor comprising an electrolytic solution which comprises a solution containing a solvent consisting of 20 to 80% by weight of an organic solvent and 80 to 20% by weight of water, and

at least one electrolyte selected from the group consisting of a carboxylic acid or a salt thereof and an inorganic acid or a salt thereof, having added thereto at least one nitro compound selected from the group consisting of nitrophenol, nitrobenzoic acid, dinitrobenzoic acid, nitroacetophenone and nitroanisole.

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- 9. The electrolytic capacitor according to claim 8, wherein the nitro compound is a combination of two or more nitro compounds.
- 10. The electrolytic capacitor according to claim 8 or 9, wherein the nitro compound is added in the amount of 0.01 to 5% by weight based on the total amount of the electrolytic solution.
- 11. The electrolytic capacitor according to any one of claims 8 to 10, comprising a capacitor element formed of an anode foil, a cathode foil opposed to the anode foil and a separator disposed between the anode foil and the cathode foil, and the electrolytic solution.

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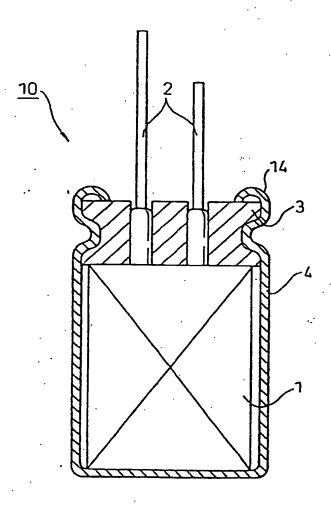
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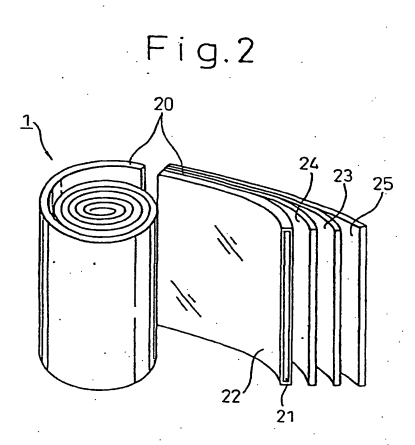
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Fig.1





INTERNATIONAL SEARCH REPORT

International application No.

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Category*	Citation of document, with indication, where	appropriate, of the relevan	nt passages	Relevant to claim No.	
•	JP, 5-205978, A (Nichikon Cor 13 August, 1993 (13.08.93)	poration), (Family: none)		1-6,8-11	
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Y JP, 1-168017, A (Nichikon Corporation), 03 July, 1989 (03.07.89) (Family: none)					
03 July, 1989 (03.07.89) (Family: none)					
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	documents are listed in the continuation of Box C.	See patent family			
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